

SECURAL SECURIOR SECURES SECURES MANAGER

MICROCOPY RESOLUTION TEST CHART

SOCI MERCESSES SOCIONES PROPERTO



REPORT DOCUMENTATION PAGE	READ INSTRUCTIONS BEFORE COMPLETING FORM		
REPORT NUMBER 2. GOVT ACCESSION NO.	3. RECIPIENT'S CATALOG NUMBER		
Organophosphazenes. 19. Copolymerization of 2-(2-Ethoxyvinyl)pentafluorocyclotriphosphazene	5. туре оf пероят & релюд соvene Technical Report		
with Styrene and Methyl Methacrylate	6. PERFORMING ORG. REPORT NUMBER		
AUTHOR(a)	B. CONTHACT OR GRANT NUMBER(s)		
Christopher W. Allen and Randall P. Bright	N00014-85-K		
PERFORMING ORGANIZATION NAME AND ADDRESS Department of Chemistry University of Vermont Burlington, Vermont 05405	10. PROGRAM ELEMENT, PROJECT, TASK AREA & WORK UNIT NUMBERS		
CONTROLLING OFFICE NAME AND ADDRESS Department of the Navy	12. REPORT DATE		
Office of Naval Research	3/14/86		
	13. NUMBER OF PAGES		
Arlington, VA 22217			
	15. SECURITY CLASS. (of this report)		
Arlington, VA 22217	15. SECURITY CLASS. (of this report) unclassified		

17. DISTRIBUTION STATEMENT (of the abstract entered in Block 20, if different from Report)

SELECTE MAR 2 5 1986

18. SUPPLEMENTARY NOTES

Accepted for publication in macromolecules

19. KEY WORDS (Continue on reverse side if necessary and identify by block number) cyclophosphazenes copolymerization reactivity ratios flame retardant

20. ABSTRACT (Continue on reverse side if necessary and identify by block number)

The novel cyclophosphazene, 2-(2-ethoxyvinyl)pantafluorocyclotriphosphazene (NypyF5C(OC)H5)=CH2) undergoes radical copolymerization with styrene and methyl methacrylate. The styrene system was examined in detail with flame retardant copolymers having up to 43.7% phosphazene content being obtained. Reactivity ratios for the styrene-ethoxyvinylphosphazene copolymerization have been calculated by several methods. An examination of the Alfrey-Price parameters for the phosphazene indicates that the major pertubation of the olefinic center is (continued on reverse, block 22a)

DD 1 JAN 73 1473 EDITION OF 1 NOV 65 IS OBSOLETE

22a.

100HB \$10

through the o electron withdrawing effect of the phosphazene. The copolymer molecular weights decrease with increasing phosphazene content. The thermal decomposition of the copolymers is a two step process with the phosphazene being involved in the first step.

OFFICE OF NAVAL RESEARCH
Contract NO0014-85-K-0558
Task No. NR413C012+01
TECHNICAL REPORT NO. 1

Organophosphazenes. 19. Copolymerization of 2-(2-Ethoxyviny1)pentafluorocyclotriphosphazene with Styrene and Methyl Methacrylate

bу

Christopher W. Allen and Randall P. Bright

Accepted for Publication
in
Macromolecules

University of Vermont Department of Chemistry Burlington, VT 05405

March 14, 1986

Reproduction in whole or in part is permitted for any purpose of the United States Government

This document has been approved for public release and sale; its distribution is unlimited

Organophosphazenes. 19. Copolymerization of 2-(2-Ethoxyvinyl)pentafluorocyclotriphosphazene with Styrene and Methyl Methacrylate¹

Christopher W. Allen* and Randall P. Bright

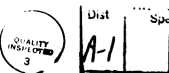
Department of Chemistry, University of Vermont
Burlington, Vermont 05405. Received

ABSTRACT: The novel cyclophosphazene, 2-(2-ethoxyvinyl)pentafluorocyclotriphosphazene ($N_3P_3F_5C(0C_2H_5)$ =CH₂) undergoes radical copolymerization with styrene and methyl methacrylate. The styrene system was examined in detail with flame retardant copolymers having up to 43.7% phosphazene content being obtained. Reactivity ratios for the styrene-ethoxyvinylphosphazene copolymerization have been calculated by several methods. An examination of the Alfrey-Price parameters for the phosphazene indicates that the major pertubation of the olefinic center is through the σ electron withdrawing effect of the phosphazene. The copolymer molecular weights decrease with increasing phosphazene content. The thermal decomposition of the copolymers is a two step process with the phosphazene being involved in the first step.

Introduction

44 - - - 3555555 3555555 42226564

Recent work in our laboratories has been devoted to the synthesis and reactions of organofunctional phosphazenes. Of particular interest is the preparation of polymeric systems derived from organofunctional phosphazene monomers. We originally demonstrated the feasibility of such an approach by reporting the copolymerization of 2-(2-propenyl)pentafluorocyclotriphosphazene,



 $N_3P_3F_5C(CH_3)=CH_2$, with styrene and vinyl benzyl chloride.³ These copolymers met our initial criteria of being hybrid organic-inorganic polymers, exhibiting significant flame retardant behavior³ and having a functionalized surface due to the presence of the halophosphazene.⁴ There are, however, problems with the inorganic monomer which are related to the polarity induced in the olefin by the highly electron withdrawing⁵⁻⁷ phosphazene substituent. The olefin polarity causes difficulties in the preparation of the monomer^{8,9} and favors termination in copolymerization reactions.³ We have recently reported the preparation of a new alkenylphosphazene monomer system derived from vinylethers in which it was proposed that the electron donor effect of the alkoxy group towards the olefin would counterbalance the electron withdrawing effect of the phosphazene.^{9,10} This expectation was realized in the synthesis of the monomer. This paper reports the copolymerization behavior of the 2-(2-ethoxyvinyl)pentafluorocyclotriphosphazene, $N_3P_3F_5C(OC_2H_5)=CH_2$, monomer.

Experimental

Materials. Hexachlorocyclotriphosphazene (Ethyl Corp.) was converted to hexafluorocyclotriphosphazene 11 which in turn was converted to (2-ethoxy-vinyl)pentafluorocyclotriphosphazene 10 by previously reported procedures. Styrene (Aldrich) was dried over P_2O_5 , distilled and stored in a sealed, dark bottle over nitrogen at $O^{\circ}C$. Prior to use, a small amount of styrene was added to methanol. If the solution remained clear, it was assumed that no spontaneous polymerization had occurred. Methyl methacrylate (Aldrich) was purified in a manner identical to that used for styrene. Azobisisobutyronitrile, AIBN, (Aldrich) was recrystallized from ethanol/water. All other reagents and solvents were obtained from standard sources and used as received.

Measurements. Infrared spectra were obtained as KBr pellets using a Nicolet

6000 FT-IR spectrophotometer. Gel permeation chromatography was performed on a Waters A.V.C. 202 high pressure liquid chromatograph equipped with Waters 10^3\AA and 10^4\AA microstyragel columns using toluene as the eluting solvent. The columns were calibrated with polystyrene standards (Waters). Thermal analyses were recorded on a DuPont 900/950 Thermal Analyzer with the sample in He flow atmosphere and a 10^6C/min heating rate. Membrane osmometric data was obtained on WESCAN Model 230 Recording Membrane Osmometer using toluene as the solvent. Data at four concentrations (1.02 to 3.36 g/l) were extrapolated to infinite dilution. Vapor pressure osmometry was performed by Schwartzkopf Microanalytical Laboratory. Elemental analyses were performed by Integral Microanalytical Laboratories. Reactivity ratios were calculated using both linear (Fineman-Ross¹² and Kelen-Tudos¹³) methods with least squares best fits of the data and the Mortimer-Tidwell non-linear least squares approach. 14

KCCCCCC

Copolymerization of (2-ethoxyvinyl)pentafluorocyclotriphosphazene and styrene. Styrene and freshly distilled (25-30 $^{\circ}$ C; 0.02 mmHg) N₃P₃F₅C(0C₂H₅)=CH₂ in various mole ratios were placed in a thick walled reaction tube along with 2% AIBN. The reaction tube was sealed with a septum and flushed with nitrogen for 15 minutes through an exit needle. The tube was then placed in a constant temperature bath (57 $^{\circ}$ C \pm 1 $^{\circ}$ C) for 18 hours. Several polymerizations to be used in reactivity ratio calculations were run limiting the % conversion. The polymerization times for these reactions were thus decreased to between 2 and 4 hours. At the end of this time period, the reaction was halted by opening the reaction vessel, and adding 5-10 ml. methylene chloride. The solution was filtered to remove any insoluble material, and then added dropwise to a stirred solution of 250 ml. methanol. The copolymer precipitates as a white powdery product. After filtration, the polymer was dissolved in methylene chloride, reprecipitated and

dried in vacuo. The composition yield and molecular weight data for these copolymers are given in Table I.

Copolymerization of (2-ethoxyvinyl)pentafluorocyclotriphosphazene and methyl methacrylate.

The methyl methacrylate (MMA) was purified in an identical manner to styrene. Reaction conditions and workup procedures utilized were identical to that described for the styrene copolymerizations. An equal molar mixture of MMA and $N_3P_3F_5C(OC_2H_5) = CH_2$ was polymerized to 6.77% conversion, and produced a copolymer which had a composition (MMA/ $N_3P_3F_5C(OC_2H_5) = CH_2$) of 3.15.

Attempted Homopolymerization of (2-ethoxyvinyl)pentafluorocyclotriphosphazene.

Numerous attempts were made at homopolymerization of $N_3P_3F_5C(0C_2H_5) = CH_2$ using radical (AIBN, benzoyl peroxide), redox $((NH_4)_2Fe(SO_4)_2 \cdot 6H_2O/Na_4P_2O_7 \cdot 10H_2O^{15})$, anionic (KHCO₃, t-C₄H₉Li) and cationic [BF₃ · O(C₂H₅)₂, H₂SO₄] initiators. The radical systems appeared to give trace amounts of oligomers while all other catalysts did not effect any polymerization.

Results and Discussion

As was the case with (2-propenyl)pentafluorocyclotriphosphazene³, homopolymerization of (2-ethoxyvinyl)pentafluorocyclophosphazene (I) was not observed. This may be ascribed to a combination of the reluctance of 1,1-disubstituted olefins to undergo homopolymerization and the strong electron withdrawing nature of the $N_3P_3F_5$ moiety.^{5,6} In principle, the electron withdrawing substituent should favor anionic initiation, however, the anionic catalyst may be captured at the P(V) site rather than at the olefinic center.

Copolymerization reactions involving I procede readily and the system

styrene-I was examined in detail. Infrared data show the retention of aromatic

$$N_{3}P_{3}F_{5}C = CH_{2} + \times CH = CH_{2} \xrightarrow{\Delta} \begin{bmatrix} (CH - CH_{2})(C - CH_{2})y \end{bmatrix} \cap X = 0.41 \text{ to } 7.22$$
AIBN
$$C_{6}H_{5} \xrightarrow{\Delta} \begin{bmatrix} (CH - CH_{2})(C - CH_{2})y \end{bmatrix} \cap Y = 0.04 \text{ to } 0.77$$

TOO YESTEN STATES

and phosphazene bonds and the absence of all olefinic absorptions in the copolymers. The compositional, conversion and molecular weight data for the new copolymers are shown in Table I. It should be noted that reactions 1-6 were stopped at low conversions. This was necessary for certain of the calculations of reactivity ratios. The copolymers are white, powdery substances, which dissolve in methylene chloride, toluene, benzene, tetrahydrofuran, diethyl ether, and acetone. They are stable to air, and dilute acid or base. The copolymers were also found to be flame retardant in simple flame tests.

The styrene-I copolymers may be compared to the previously reported styrene- $N_3P_3F_5C(CH_3)=CH_2$ system. The maximum phosphazene incorporation in the propenylphosphazene series was 36.7 mole % which was achieved with feed ratio (styrene phosphazene) of 4.1. The maximum incorporation of I is 43.7 mole % which was achieved with a 2.44.1 feed ratio. Feed ratios which are higher in I do not undergo polymerization. The higher incorporation of I compared to the propenyl analog indicates a significant difference in the vinylphosphazene reactivity in these two systems and is consistent with our proposal of reduction of olefin polarity by a counterbalancing of the electron withdrawing effect of the phosphazene with the donor effect of the alkoxy group.

In order to provide a more thorough examination of the reactivity of I, we have calculated the reactivity ratios $(r_1 \text{ and } r_2)$ for the styrene-I system. These data, calculated by three different methods, may be found in Table II.

In comparing the three methods, it is observed that all give similar results with the results of both of the linearization methods falling within the 95% joint confidence limits of the Mortimer-Tidwell calculation. Since it is expected that the Mortimer-Tidwell approach will give the most reliable data 14, those values of \mathbf{r}_1 and \mathbf{r}_2 were used to calculate the Alfrey-Price parameters 16 with values of Q=0.18 and e=-0.06 being obtained. It is apparent upon comparison of the reactivity ratios of styrene (M_1) -ethylvinyl ether (M_2) $(r_1=80, r_2=0)^{17}$ to those of the styrene-I system that the olefinic center in I does not behave as a typical vinylether. The addition of a bulky group such as the phosphazene ring to the vinylether might be expected to offer some steric inhibition to copolymerization so the observed increase in activity indicates that the electronic structure of the vinylether is significantly perturbed by the $N_3P_3F_5$ moiety. Similar conclusions were drawn from the nmr data on I. 10 Some insight into the mechanism of modification of the olefin electronic structure can be gained from a comparison of the Q and e values for I to those for related monomers. 17 The Q value of I lies in the general range of those observed for vinyl ethers thus indicating no major mesomeric interaction between the phosphazene and the olefin. The reluctance of the phosphazene unit to enter into significant conjugative interactions with unsaturated organic moieties has been previously noted. 6 The approximate nature of the Q, e data does not prohibit small amounts of mesomeric interactions occuring in I such as has been claimed in other organophosphazenes. 7,18 The e value for I is quite different from those typically found for vinylethers and this difference may be ascribed to the reduction of the electron rich nature of the vinylether by the strong σ electron withdrawing effect of the phosphazene moiety.

THE PROPERTY OF THE PARTY OF TH

If the Q and e values for I are combined with the literature 17 values for methyl methacrylate (MMA), the reactivity ratios for the MMA-I copolymerization

may be calculated. For an equimolar copolymerization of MMA and I, the copolymer composition calculated from the reactivity ratios and the copolymer equation is $(\text{MMA})_{3.5}(I)_1$ which is in reasonable agreement (given the nature of the data) with the observed composition of $(\text{MMA})_{3.15}(I)_1$.

The reactivity ratios of the styrene-I system show that a styryl radical has a greater tendency to add to styrene than to I. Using a value of 165 l/mole-sec for the styrene homopropagation rate constant (k_{11}) , 19 we can estimate (from r_1) a value of k_{12} of 54.3 l/mole-sec, i.e. roughly a three-fold preference for styrene homo vs. cross propagation. The reluctance of I to undergo homopolymerization, suggests that the homopropagation rate constant of I, k_{22} , is diminishingly small. To get a value of r_2 of 0.19, the rate constant for the addition of the radical of I to styrene (k_{21}) must be large. The combination of large values of k_{11} and k_{21} lead to copolymers which are rich in styrene.

Molecular weight data for the styrene-I copolymers may be found in Table I. The membrane osmometry data is significantly higher than the gel permeation chromatography (gpc) data. If the gpc data is close to the true value, then the osmometric data would be artificially high due to diffusion of low molecular weight fragments across the membrane. Alternatively, polystyrene may not be a good gpc calibrant for the styrene-I copolymers. A low molecular weight polymer was selected for study by vapor phase osmometry and it was found that the molecular weight was higher than the gpc value in roughly the same ratio as the membrane osmometry to gpc data. It appears that the gpc value are uniformly low and that the true values are closer to those obtained by the absolute methods. One possible explanation for the low gpc values would be electrostatic attraction between the phosphazene and phenyl rings resulting in a reduction in size of the copolymer over the expected average dimensions of an unperturbed chain. The general trend in molecular weights is similar to that observed for the

styrene-propenylphosphazene copolymers, i.e. a reduction in molecular weight with increased phosphazene content. The high reactivity and low homopropogation rate of the phosphazene radical would result in increased occurance of termination steps with increased amounts of phosphazene in the system.

The TGA data for thermal decomposition of the styrene-I copolymers are found in Table III. In each case, the decomposition is, at least, a two step process with a typical TGA scan shown in Figure 1. There is a correlation between the % weight loss in the first step and the weight % of I in the copolymer. Given this observation, it is reasonable to assume that the first stage of decomposition involves the ethoxyvinylphosphazene region of the copolymer. There is also a lowering of the onset temperature for decomposition and the temperature required for 50% weight loss with increasing phosphazene content. These observations also suggest that the incorporation of I contributes to the destabilization of the copolymer to thermal decomposition. It is of interest to note the lower thermal stability of the ethoxyvinyl- compared to propenylphosphazene³ copolymers.

In summary, we have developed a new series of novel copolymers involving a vinyl phosphazene and traditional organic monomers. Significant alterations of copolymerization behavior of vinylphosphazene monomers may be effected by the nature of the substituent on the olefinic center. The nature of the effect of the phosphazene on the vinylether center in I has been elucidated by a consideration of the Q and e values for this system.

Acknowledgements. We wish to thank Dr. G.A. Stahl of Phillips Petroleum Co. for a copy of the Mortimer-Tidwell program and Mr. D.E. Brown for adapting it to our computer. This work was supported, in part, by the Office of Naval Research.

References

- (1) Part 18. Allen, C.W.; Bedell, S.; Cordes, W.A.; Pennington, W.T. <u>Inorg.</u>
 Chem. 1985, 24, 1653.
- (2) Allen, C.W. <u>J. Polym. Sci. Polym. Symposia</u> 1983, <u>70</u>, 79.
- (3) Dupont, J.G.; Allen, C.W. Macromolecules 1979, 12, 169.
- (4) Allen, C.W.; Dupont, J.G. Ind. Eng. Chem. Prod. Res. Dev. 1979, 18, 80.
- (5) Allen, E.W. J. Organometallic Chem. 1977, 125, 215.
- (6) Allen, C.W.; Green, J.C. Inorg. Chem. 1980, 19, 1719.
- (7) Harris, P.J., Williams, K.B., Fisher, B.L. J. Org. Chem. 1984, 49, 406.
- (8) Dupont, J.G., Allen, C.W. <u>Inorg. Chem.</u> 1978, <u>17</u>, 3093.
- (9) Allen, C.W., Bright, R.P., Ramachandran, K. "ACS Symp. Ser. No. 171", Quinn, L.D., Verkade, J.G., Eds., Amer. Chem. Soc.: Washington, D.C., 1981, p. 321.
- (10) Allen, C.W., Bright, R.P. <u>Inorq. Chem</u> 1983, <u>22</u>, 1291.
- (11) Moeller, T., John, K., Tsang, F. Chem. Ind. (London) 1961, 347.
- (12) Fineman, M.; Ross, S. <u>J Poly</u>. <u>Sci</u>. **1950**, <u>5</u>, 259.
- (13) Tüdös, F.; Kelen, T.; Földes-Berenzwich, F.; Turcsanyi, B. J. Macromol.
 Sci. Chem., 1976, A10, 1513. (b) Tüdös, F.; Kelen, T. Ibid, 1975, A9, 1.
 (c) Tüdös, F.; Kelen, T. Reaction Kin. Cat. Letters, 1974, 1, 487.
- (14) Tidwell, P.W., Mortimer, G.A. J. Poly. Sci. A-1, 1965, 3, 369. (b)
 Tidwell, P.W., Mortimer, G.A. J. Macromol. Sci. Macromol. Chem. Rev.,
 1970, 4, 281.
- (15) Marvel, C.S., Deanin, R., Overberger, C.G., Kuhn, B.M. <u>J. Poly. Sci.</u>, 1948, 3, 128.
- (16) Alfrey Jr., T.; Young, L.J. "Copolymerization", 2nd ed; Ham, G.E., Ed.; Wiley; New York, 1971.

- (17) Young, L.J. **Polymer Handbok**, 2nd ed; Brandrup, J., Immergut, E.H., Eds.; Wiley: New York, 1975; Chapter II.
- (18) Krishnamurthy, S.S.; Ramabraman, P.; Woods, M. <u>Org. Magn. Reson.</u>, 1981, <u>15</u>, 205.

basis become alabatic bulleting

(19) Odian, G. **Principles of Polymerization**, 2nd Ed; Wiley: New York, 1981, p. 258.

Composition, Yield and Molecular Weight Data^a for Styrene(II)-Ethoxyvinylpentafluoro-Table I

cyclotriphosphazene(I) Copolymers

	_1	_											_		
1:	E _S	6.67		7.15	3.83	8.79	•	2.18	2.48	2.91	2.55	1.36	1.74		
	Σ	70.7K		71.2K	33.4K	56.1K	1	15.0K	7.8K	6.6K	4.2K	3.3K	6.5K		
	IΣC	10.6K 36.1K	9.8K 31.0K ^C	10.0K	8.7K	6.4K	ı	6.9K	3.2K	2.3K	1.6K 5.9Kd	2.4K	3.8K		
	% Conversion (Wt)	11.4	12.0	9.4	8.0	7.5	1.8	61.3	55.4	52.7	35.0	22.4	18.0	0.0	
•	Reaction Time (hrs)	7	2	2	8	8	4	18	18	18	18	18	18	18	
	Product Ratio (I/II)	24.22	20.64	15.69	13.64	9.11	3.62	5.78	4.20	3.94	2.80	2.06	1.30	1	
	Feed Ratio (I/II)	7.22	5.78	4.82	3.85	2.89	66.	2.94	2.02	2.00	1.00	.80	.41	.25	
	Run	-	2	٣	4	۲,	9	7	ထ	٥	10	11	12	13	•

 $^{^{}a}$ Molecular weights measured by gel permeating chromatography, except where indicated. As determined from weight percent nitrogen. Smembrane Osmometry.

demorane Usmometry.

dvapor Pressure Osmometry.

Table II

Reactivity Ratios for Styrene-Ethoxyvinylpentafluorocyclotriphosphazene Copolymerization Reaction

Method	$\frac{\mathbf{r_l}^a}{\mathbf{l}}$	$\frac{\mathbf{r_2}^{\mathbf{b}}}{2}$
Fineman-Ross	3.40	0.30
Kelen-Tudos	3.56	0.28
Mortimer-Tidwell	3.04	0.19

a
$$M_1$$
=Styrene b M_2 = N_3 P_3 F_5 $C(0C_2$ $H_5)=CH_2$

THE PROPERTY OF SECOND STATES OF THE PARTY O

Table III

TGA Data for Styrene-Ethoxyvinylpentafluorocyclotriphosphazene Copolymers

Wt % Phosphazene in Copolymer	Weight loss range	% Wt. loss first step	T ₅₀ (°C)
33.4	205-280	24.4	350
40.8	200-280	30.3	350
42.5	200-290	28.3	355
50.8	190-270	39.3	320
69.2	180-260	46.7	315

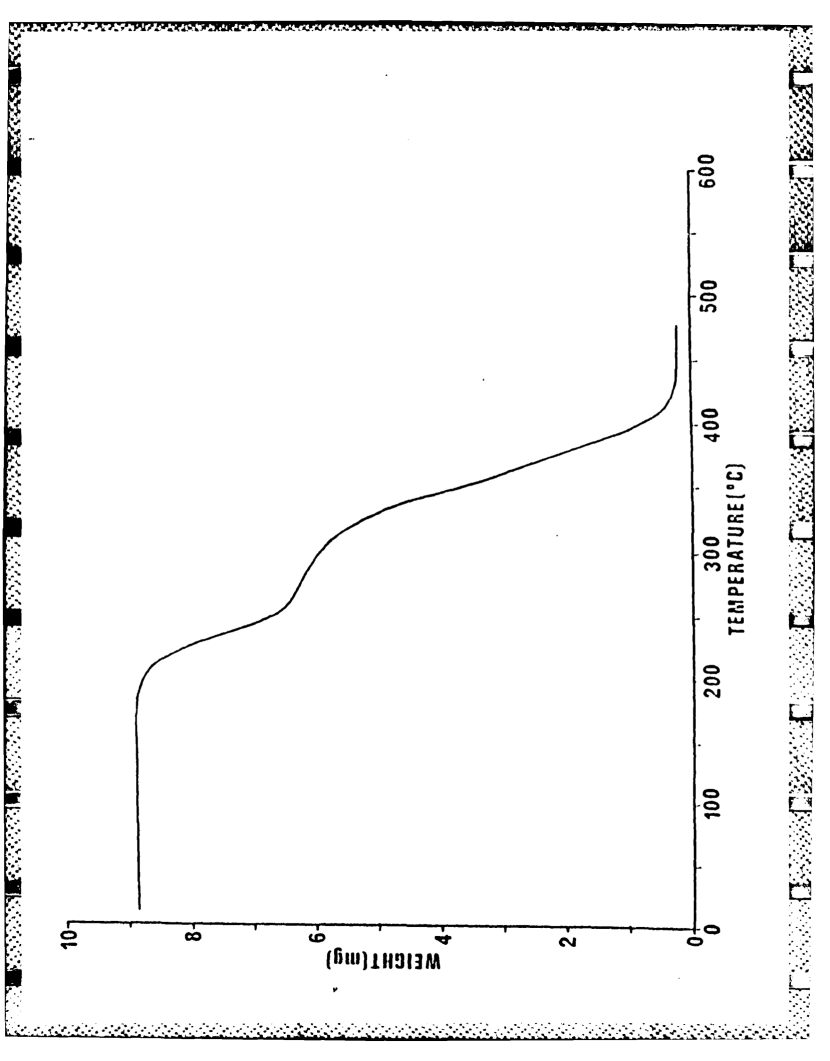


Figure 1. A Typical TGA Scan for Styrene-Ethoxyvinylpentafluorocyclotriphosphazene Copolymers. 14

CONTRACTOR CONTRACTOR

-IIMED 1-86